OECD GUIDELINE FOR THE TESTING OF CHEMICALS

DRAFT PROPOSAL FOR A NEW TEST GUIDELINE

Skin Sensitisation: Local Lymph Node Assay: BrdU-ELISA

INTRODUCTION

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- 5 1. OECD Guidelines for the Testing of Chemicals are periodically reviewed in light of
- 6 scientific progress, changing regulatory needs, and animal welfare considerations. The first Test Guideline (TG) for the determination of skin sensitisation in the mouse, the 7
- 8 radiolabelled Local Lymph Node Assay (LLNA; TG 429) was adopted in 2002 (1). The
- 9
- details of the validation of the LLNA and a review of the associated work have been 10 published (2)(3)(4)(5)(6). A modified Local Lymph Node Assay (LLNA) utilising non-
- radiolabelled 5-bromo-2-deoxyuridine (BrdU) (Chemical Abstracts Service [CAS] No 59-14-11
- 12 3) ELISA test method (LLNA: BrdU-ELISA) has been validated and based on a formal peer
- review, the LLNA: BrdU-ELISA is considered useful for identifying skin sensitising and 13
- 14 non-sensitising test substances, with certain limitations (7)(8)(9). This is the fourth Test
- Guideline to be promulgated for assessing skin sensitisation potential of chemicals in 15
- animals. Test Guideline 406 utilises guinea pig tests, notably the guinea pig maximisation 16
- 17 test and the Buehler test (10). The LLNA: BrdU-ELISA provides certain advantages over TG
- 18 406 with regard to both scientific progress and animal welfare.
- 19 2. Similar to the LLNA, the LLNA: BrdU-ELISA studies the induction phase of skin
- 20 sensitisation and provides quantitative data suitable for dose response assessment.
- 21 Furthermore, an ability to detect skin sensitizers without the necessity for using a radiolabel
- 22 for DNA eliminates the potential for occupational exposure to radioactivity and waste
- 23 disposal issues. This in turn may allow for the increased use of mice to detect skin 24 sensitizers, which could further reduce the use of guinea pigs to test for skin sensitisation
- 25 potential (i.e. TG 406) (10). A reduced LLNA: BrdU-ELISA (rLLNA: BrdU-ELISA)
- protocol that uses fewer animals is also described in this TG (1)(12)(13). The rLLNA: BrdU-26
- 27 ELISA may be used for the hazard classification of skin sensitising test substances when
- dose-response information will not be needed provided there is adherence to all other 28
- 29 LLNA: BrdU-ELISA protocol specifications as described in this TG. The rLLNA: BrdU-
- 30 ELISA should not be used for the hazard identification of skin sensitising test substances
- 31 when dose-response information is needed.

DEFINITIONS

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33 3. Definitions used are provided in Annex 1.

34 INITIAL CONSIDERATIONS AND LIMITATIONS

- 35 4. The LLNA: BrdU-ELISA is a modified LLNA method for identifying potential skin
- sensitising test substances, with specific limitations. This does not necessarily imply that in 36
- 37 all instances the LLNA: BrdU-ELISA should be used in place of the LLNA or guinea pig

- tests (*i.e.* TG 406) (10), but rather that the assay may be employed as an alternative in which positive and negative results generally no longer require further confirmation (7)(8). The testing laboratory should consider all available information on the test substance prior to conducting the study. Such information will include the identity and chemical structure of the test substance; its physicochemical properties; the results of any other *in vitro* or *in vivo* toxicity tests on the test substance; toxicological data on structurally related test substances.
 - 5. The LLNA: BrdU-ELISA is an *in vivo* method and, as a consequence, will not eliminate the use of animals in the assessment of allergic contact sensitising activity. It has, however, the potential to reduce the number of animals required for this purpose (e.g. reducing the number of guinea pigs used when the LLNA: BrdU-ELISA is used instead of guinea pig assays and the LLNA where the use of radioactivity is discouraged). Moreover, the LLNA: BrdU-ELISA offers a substantial refinement of the way in which animals are used for allergic contact sensitisation testing. The LLNA: BrdU-ELISA is based upon consideration of immunological events stimulated by chemicals during the induction phase of sensitisation. Unlike guinea pig tests (i.e. TG 406) (10), the LLNA: BrdU-ELISA does not require that challenge-induced dermal hypersensitivity reactions be elicited. Furthermore, the LLNA: BrdU-ELISA does not require the use of an adjuvant, as is the case for the guinea pig maximisation test, as described in reference (10). Thus, the LLNA: BrdU-ELISA reduces animal distress. Despite the advantages of the LLNA: BrdU-ELISA over TG 406 (10), there are certain limitations that may necessitate the use of TG 406 (10) (e.g. the testing of certain metals, false positive findings with certain skin irritants [such as some surfactant-type materials] (6)(14), solubility of the test material, or test substance classes or materials containing functional groups shown to act as potential confounders (15). Limitations that have been identified for the LLNA have been recommended to apply also to the LLNA: BrdU-ELISA (7). For the validation database of 43 substances, the LLNA: BrdU-ELISA correctly identified all 32 LLNA sensitizers, but two of 11 LLNA non-sensitizers were identified as borderline positive, with Stimulation Index (SI) values between 1.6 and 1.9 (7). Other than such identified limitations and considerations, the LLNA: BrdU-ELISA should be applicable for testing any test substances unless there are properties associated with these materials that may interfere with the accuracy of the LLNA: BrdU-ELISA.

PRINCIPLE OF THE TEST

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6. The basic principle underlying the LLNA: BrdU-ELISA is that sensitizers induce proliferation of lymphocytes in the lymph nodes draining the site of test substance application. This proliferation is proportional to the dose and to the potency of the applied allergen and provides a simple means of obtaining a quantitative measurement of sensitisation. Proliferation is measured by comparing the mean proliferation in each test group to the mean proliferation in the vehicle treated control group. The ratio of the mean proliferation in each treated group to that in the concurrent vehicle treated control group, termed the SI, is determined, and should be ≥1.6 before further evaluation of the test substance as a potential skin sensitizer is warranted. The methods described here are based on the use of measuring BrdU content to indicate an increased number of proliferating cells in the draining auricular lymph nodes. BrdU is an analogue of thymidine and is similarly incorporated into the DNA of proliferating cells. The incorporation of BrdU is measured by ELISA, which utilises an antibody specific for BrdU that is also labelled with peroxidase.

- When the substrate is added, the peroxidase reacts with the substrate to produce a coloured
- product that is quantified at a specific absorbance using a microtiter plate reader.

84 <u>DESCRIPTION OF THE ASSAY</u>

Selection of animal species

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- 86 7. The mouse is the species of choice for this test. Validation studies for the
- 87 LLNA: BrdU-ELISA were conducted exclusively with the CBA/JN strain, which is therefore
- 88 considered the preferred strain (7)(9). Young adult female mice, which are nulliparous and
- 89 non-pregnant, are used. At the start of the study, animals should be between 8-12 weeks old,
- and the weight variation of the animals should be minimal and not exceed 20% of the mean
- 91 weight. Alternatively, other strains and males may be used when sufficient data are generated
- 92 to demonstrate that significant strain and/or gender-specific differences in the LLNA: BrdU-
- 93 ELISA response do not exist.

Housing and feeding conditions

- 95 8. Mice should be group housed (16), unless adequate scientific rationale for housing
- 96 mice individually is provided. The temperature of the experimental animal room should be
- 97 22°C (± 3°C). Although the relative humidity should be at least 30% and preferably not
- exceed 70%, other than during room cleaning, the aim should be 50-60%. Lighting should be
- 99 artificial, the sequence being 12 hours light, 12 hours dark. For feeding, conventional
- laboratory diets may be used with an unlimited supply of drinking water.

101 **Preparation of animals**

- 102 9. The animals are randomly selected, marked to permit individual identification (but
- not by any form of ear marking), and kept in their cages for at least five days prior to the start
- of dosing to allow for acclimatisation to the laboratory conditions. Prior to the start of
- treatment all animals are examined to ensure that they have no observable skin lesions.

Preparation of dosing solutions

- 107 10. Solid test substances should be dissolved or suspended in solvents/vehicles and
- diluted, if appropriate, prior to application to an ear of the mice. Liquid test substances may
- be applied neat or diluted prior to dosing. Insoluble materials, such as those generally seen in
- medical devices, should be subjected to an exaggerated extraction in an appropriate solvent
- to reveal all extractable constituents for testing prior to application to an ear of the mice. The
- test substances should be prepared daily unless stability data demonstrate the acceptability of
- storage.

114 **Reliability check**

- 115 11. Positive controls (PC) are used to demonstrate appropriate performance of the assay
- by responding with adequate and reproducible sensitivity to a sensitising test substance for
- which the magnitude of the response is well characterised. Inclusion of a concurrent PC is
- recommended because it demonstrates competency of the laboratory to successfully conduct

- each assay and allows for an assessment of intra- and inter-laboratory reproducibility and
- comparability. A PC for each study is required by some regulatory authorities. Accordingly,
- the routine use of a concurrent PC is encouraged to avoid the need for additional animal
- testing to meet such requirements that might arise from the use of a periodic PC (see
- paragraph 12). The PC should produce a positive LLNA: BrdU-ELISA response at an
- exposure level expected to give an increase in the $SI \ge 1.6$ over the negative control (NC)
- group. The PC dose should be chosen such that the induction is reproducible but not
- excessive (e.g. SI > 14 would be considered excessive). Preferred PC test substances are 25%
- 127 hexyl cinnamic aldehyde (CAS No 101-86-0) and eugenol (CAS No 97-53-0) in
- acetone: olive oil. There may be circumstances in which, given adequate justification, other
- 129 PC test substances, meeting the above criteria, may be used.
- 130 12. While inclusion of a concurrent PC group is recommended, there may be situations
- in which periodic testing (i.e. at intervals ≤ 6 months) of the PC test substance may be
- adequate for laboratories that conduct the LLNA: BrdU-ELISA regularly (i.e. conduct the
- LLNA: BrdU-ELISA at a frequency of no less than once per month) and have an established
- historical PC database that demonstrates the laboratory's ability to obtain reproducible and
- accurate results with PCs. Adequate proficiency with the LLNA: BrdU-ELISA can be
- successfully demonstrated by generating consistent positive results with the PC in at least 10
- independent tests conducted within a reasonable period of time (*i.e.* less than one year).
- 138 13. A concurrent PC group should always be included when there is a procedural
- change to the LLNA: BrdU-ELISA (e.g. change in trained personnel, change in test method
- materials and/or reagents, change in test method equipment, change in source of test
- animals), and such changes should be documented in laboratory reports. Consideration
- should be given to the impact of these changes on the adequacy of the previously established
- historical database in determining the necessity for establishing a new historical database to
- document consistency in the PC results.
- 145 14. Investigators should be aware that the decision to conduct a PC on a periodic basis
- instead of concurrently has ramifications on the adequacy and acceptability of negative study
- results generated without a concurrent PC during the interval between each periodic PC
- study. For example, if a false negative result is obtained in the periodic PC study, negative
- 149 test substance results obtained in the interval between the last acceptable periodic PC study
- and the unacceptable periodic PC study may be questioned. Implications of these outcomes
- should be carefully considered when determining whether to include concurrent PCs or to
- only conduct periodic PCs. Consideration should also be given to using fewer animals in the
- 153 concurrent PC group when this is scientifically justified and if the laboratory demonstrates,
- based on laboratory-specific historical data, that fewer mice can be used (17).
- 155 15. Although the PC test substance should be tested in the vehicle that is known to elicit
- a consistent response (e.g. acetone: olive oil), there may be certain regulatory situations in
- which testing in a non-standard vehicle (clinically/chemically relevant formulation) will also
- be necessary (1). If the concurrent PC test substance is tested in a different vehicle than the
- test substance, then a separate vehicle control for the concurrent PC should be included.

- 160 16. In instances where test substances of a specific chemical class or range of responses
- are being evaluated, benchmark test substances may also be useful to demonstrate that the
- test method is functioning properly for detecting the skin sensitisation potential of these types
- of test substances. Appropriate benchmark test substances should have the following
- 164 properties:

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- structural and functional similarity to the class of the test substance being tested;
- known physical/chemical characteristics;
- supporting data from the LLNA: BrdU-ELISA;
- supporting data from other animal models and/or from humans.

TEST PROCEDURE

Number of animals and dose levels

- 171 17. A minimum of four animals is used per dose group, with a minimum of three
- concentrations of the test substance, plus a concurrent NC group treated only with the vehicle
- for the test substance, and a PC (concurrent or recent, based on laboratory policy in
- considering paragraphs 11-15). Except for absence of treatment with the test substance,
- animals in the control groups should be handled and treated in a manner identical to that of
- animals in the treatment groups.
- 177 18. Dose and vehicle selection should be based on the recommendations given in the
- 178 references (2) and (19). Consecutive doses are normally selected from an appropriate
- 179 concentration series such as 100%, 50%, 25%, 10%, 5%, 2.5%, 1%, 0.5%, etc. Adequate
- scientific rationale should accompany the selection of the concentration series used. All
- existing toxicological information (e.g. acute toxicity and dermal irritation) and structural
- and physicochemical information on the test substance of interest (and/or structurally related
- test substances) should be considered, where available, in selecting the three consecutive
- concentrations so that the highest concentration maximises exposure while avoiding systemic
- toxicity and/or excessive local skin irritation (19)(20). In the absence of such information, an
- initial pre-screen test may be necessary (see paragraphs 21-1).
- 187 19. The vehicle should not interfere with or bias the test result and should be selected on
- the basis of maximising the solubility in order to obtain the highest concentration achievable
- while producing a solution/suspension suitable for application of the test substance.
- 190 Recommended vehicles are acetone: olive oil (4:1 v/v), N,N-dimethylformamide, methyl
- ethyl ketone, propylene glycol, and dimethyl sulphoxide (6) but others may be used if
- sufficient scientific rationale is provided. In certain situations it may be necessary to use a
- 193 clinically relevant solvent or the commercial formulation in which the test substance is
- marketed as an additional control. Particular care should be taken to ensure that hydrophilic
- materials are incorporated into a vehicle system, which wets the skin and does not
- immediately run off by incorporation of appropriate solubilisers (e.g. 1% Pluronic® L92).
- 197 Thus, wholly aqueous vehicles are to be avoided.

198 20. The processing of lymph nodes from individual mice allows for the assessment of 199 inter-animal variability and a statistical comparison of the difference between test substance 200 and vehicle control group measurements (see paragraph 34). In addition, evaluating the 201 possibility of reducing the number of mice in the PC group is only feasible when individual 202 animal data are collected (17). Further, some national regulatory authorities require the 203 collection of individual animal data. Regular collection of individual animal data provides an 204 animal welfare advantage by avoiding duplicate testing that would be necessary if the test 205 substance results originally collected in one manner (e.g. via pooled animal data) were to be 206 considered later by regulatory authorities with other requirements (e.g. individual animal 207 data).

Pre-screen test

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- 21. In the absence of information to determine the highest dose to be tested (see paragraph 18), a pre-screen test should be performed in order to define the appropriate dose level to test in the LLNA: BrdU-ELISA. The purpose of the pre-screen test is to provide guidance for selecting the maximum dose level to use in the main LLNA: BrdU-ELISA study, where information on the concentration that induces systemic toxicity (see paragraph 1) and/or excessive local skin irritation (see paragraph 23) is not available. The maximum
- dose level tested should be a concentration of 100% of the test substance for liquids or the
- maximum possible concentration for solids or suspensions, unless available information suggests that this concentration induces systemic toxicity and/or excessive local irritation
- after topical application in the mouse.
- 219 22. The pre-screen test is conducted under conditions identical to the main
- LLNA: BrdU-ELISA study, except there is no assessment of lymph node proliferation and fewer animals per dose group can be used. One or two animals per dose group are suggested.
- fewer animals per dose group can be used. One or two animals per dose group are suggested.
 All mice will be observed daily for any clinical signs of systemic toxicity or local irritation at
- the application site. Body weights are recorded pre-test and prior to termination (Day 6).
- Both ears of each mouse are observed for erythema and scored using Table 1 (20). Ear
- 225 thickness measurements are taken using a thickness gauge (e.g. digital micrometer or
- Peacock Dial thickness gauge) on Day 1 (pre-dose), Day 3 (approximately 48 hours after the
- first dose), and Day 6. Additionally, on Day 6, ear thickness could be determined by ear
- 228 punch weight determinations. Excessive local irritation is indicated by an erythema score ≥ 3
- and/or ear thickness of \geq 25% on any day of measurement (21)(22). The highest dose selected
- 230 for the main LLNA: BrdU-ELISA study will be the next lower dose in the pre-screen
- 231 concentration series (see paragraph 18) that does not induce systemic toxicity and/or
- 232 excessive local skin irritation.

233 **Table 1.** Erythema Scores

Observation	Score
No erythema	0
Very slight erythema (barely perceptible)	1
Well-defined erythema	2
Moderate to severe erythema	3
Severe erythema (beet redness) to eschar formation preventing grading of erythema	4

- 23. In addition to a 25% increase in ear thickness (21)(22), a statistically significant increase in ear thickness in the treated mice compared to control mice has also been used to identify irritants in the LLNA (22)(23)(24)(25)(26)(27)(28). However, while statistically significant increases can occur when ear thickness is less than 25% they have not been associated specifically with excessive irritation (25)(26)(27)(28)(29).
- 239 24. The following clinical observations may indicate systemic toxicity (30) when used 240 as part of an integrated assessment and therefore may indicate the maximum dose level to use 241 in the main LLNA: changes in nervous system function (e.g. pilo-erection, ataxia, tremors, 242 and convulsions); changes in behaviour (e.g. aggressiveness, change in grooming activity, 243 marked change in activity level); changes in respiratory patterns (i.e. changes in frequency 244 and intensity of breathing such as dyspnea, gasping, and rales), and changes in food and 245 water consumption. In addition, signs of lethargy and/or unresponsiveness and any clinical 246 signs of more than slight or momentary pain and distress, or a>5% reduction in body weight 247 from Day 1 to Day 6 and mortality should be considered in the evaluation.

Reduced LLNA

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249 25. Use of an rLLNA: BrdU-ELISA protocol instead of the multi-dose LLNA: BrdU-250 ELISA has the potential to reduce the number of animals used in a test by omitting the 251 middle and low dose groups (1)(12)(13). The reduction in number of dose groups is the only 252 difference between the LLNA: BrdU-ELISA and the rLLNA: BrdU-ELISA test method 253 protocols and for this reason the rLLNA: BrdU-ELISA does not provide dose-response 254 information. Therefore, the rLLNA: BrdU-ELISA should not be used when dose-response 255 information is needed. Like the multi-dose LLNA: BrdU-ELISA, the test substance 256 concentration evaluated in the rLLNA: BrdU-ELISA should be the maximum concentration 257 that does not induce overt systemic toxicity and/or excessive local skin irritation in the mouse 258 (see paragraph 18).

Main study experimental schedule

- 260 26. The experimental schedule of the assay is as follows:
- Day 1:
 Individually identify and record the weight of each animal and any clinical observations. Apply 25 μL of the appropriate dilution of the test substance,

- 264 the vehicle alone, or the concurrent PC (see paragraphs 11-15), to the dorsum of each ear.
 - Days 2 and 3:

Repeat the application procedure carried out on Day 1.

• Days 4:

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- No treatment.
- *Days 5:*

Inject 0.5 mL (5 mg/mouse) of BrdU (10 mg/mL) solution interperitoneally.

• *Day 6:*

Record the weight of each animal and any clinical observations. Approximately 24 hours (24 h) after BrdU injection humanely kill the animals. Excise the draining auricular lymph nodes from each mouse ear and process separately in phosphate buffered saline (PBS) for each animal. Details and diagrams of the node identification and dissection can be found in reference (17). To further monitor the local skin response in the main study, additional parameters such as scoring of ear erythema or ear thickness measurements (obtained either by using a thickness gauge, or ear punch weight determinations at necropsy) may be included into the study protocol.

Preparation of cell suspensions

- 283 27. From each mouse, a single-cell suspension of lymph node cells (LNC) excised 284 bilaterally is prepared by gentle mechanical disaggregation through 200 micron-mesh 285 stainless steel gauze or another acceptable technique for generating a single-cell suspension 286 (e.g. use of a disposable plastic pestle to crush the lymph nodes followed by passage through a #70 nylon mesh). The procedure for preparing the LNC suspension is critical in this assay 287 288 and therefore every operator should establish the skill in advance. Further, the lymph nodes 289 in negative control animals are small, so careful operation is important to avoid any artificial 290 effects on SI values. In each case, the target volume of the LNC suspension should be 291 adjusted to a determined optimised volume (approximately 15 mL). The optimised volume is 292 based on achieving a mean absorbance of the negative control group within 0.1-0.2.
- 293 <u>Determination of cellular proliferation (measurement of BrdU content in DNA of lymphocytes)</u>
- 295 28. BrdU is measured by ELISA using a commercial kit (e.g. Roche Applied Science,
- Mannheim, Germany, Catalogue Number 11 647 229 001). Briefly, 100 ul of the LNC
- suspension is added to the wells of a flat-bottom microplate in triplicate. After fixation and
- denaturation of the LNC, anti-BrdU antibody is added to each well and allowed to react.
- 299 Subsequently the anti-BrdU antibody is removed by washing and the substrate solution is
- 300 then added and allowed to produce chromogen. Absorbance at 370 nm with a reference
- 301 wavelength of 492 nm is then measured. In all cases, assay test conditions should be
- optimised (see paragraph 27).

OBSERVATIONS

Clinical observations

- 305 29. Each mouse should be carefully observed at least once daily for any clinical signs,
- 306 either of local irritation at the application site or of systemic toxicity. All observations are
- 307 systematically recorded with records being maintained for each mouse. Monitoring plans
- 308 should include criteria to promptly identify those mice exhibiting systemic toxicity, excessive
- irritation, or corrosion of skin for euthanasia.

310 **Body weights**

- 311 30. As stated in paragraph 26, individual animal body weights should be measured at
- 312 the start of the test and at the scheduled kill.

313 CALCULATION OF RESULTS

- 314 31. Results for each treatment group are expressed as the mean SI. The SI is derived by
- 315 dividing the mean BrdU labelling index/mouse within each test substance group and the
- 316 concurrent PC group by the mean BrdU labelling index for the solvent/vehicle control group.
- 317 The average SI for vehicle treated controls is then one.
- 318 The BrdU labelling index is defined as:
- 319 BrdU labelling index = $(ABS_{em} ABS blank_{em}) (ABS_{ref} ABS blank_{ref})$
- where em = emission wavelength and ref = reference wavelength.
- 321 32. The decision process regards a result as positive when $SI \ge 1.6$ (7). However, the
- 322 strength of the dose-response, the statistical significance and the consistency of the
- 323 solvent/vehicle and positive control responses may also be used when determining whether a
- borderline result is declared positive (3)(6) (1).
- 325 33. For a borderline positive response between an SI of 1.6 and 1.9, users may want to
- 326 consider additional information such as dose response, evidence of systemic toxicity or
- 327 excessive irritation, and where appropriate, statistical significance together with SI values to
- 328 confirm that such results are positives (7). Consideration should also be given to various
- 329 properties of the test substance, including whether it has a structural relationship to known
- skin sensitizers, whether it causes excessive skin irritation in the mouse, and the nature of the
- dose response seen. These and other considerations are discussed in detail elsewhere (4).
- 332 34. Collecting data at the level of the individual mouse will enable a statistical analysis
- for presence and degree of dose response in the data. Any statistical assessment could include
- 334 suitably adjusted comparisons of test groups (e.g. pair-wise dosed group versus concurrent
- 335 solvent/vehicle control comparisons). Statistical analyses may include, for instance, linear
- regression or William's test to assess dose-response trends, and Dunnett's test for pair-wise
- 337 comparisons. In choosing an appropriate method of statistical analysis, the investigator
- 338 should maintain an awareness of possible inequalities of variances and other related problems
- that may necessitate a data transformation or a non-parametric statistical analysis. In any
- case, the investigator may need to carry out SI calculations and statistical analyses with and
- without certain data points (sometimes called "outliers").

342 **DATA AND REPORTING**

- 343 **Data**
- 344 35. Data should be summarised in tabular form showing the individual animal BrdU
- labelling index values, the group mean BrdU labelling index/animal, its associated error term
- 346 (e.g. SD, SEM), and the mean SI for each dose group compared against the concurrent
- 347 solvent/vehicle control group.

348 Test report

- 349 36. The test report should contain the following information:
- 350 Test substance and control test substance:
- identification data (e.g. CAS number, if available; source; purity; known
- impurities; lot number);
- 553 physical nature and physicochemical properties (e.g. volatility, stability,
- 354 solubility);
- 355 if mixture, composition and relative percentages of components.
- 356 Solvent/vehicle:
- identification data (purity; concentration, where appropriate; volume used);
- 358 justification for choice of vehicle.
- 359 Test animals:
- 360 source of CBA mice;
- microbiological status of the animals, when known;
- of animals; number and age of animals;
- source of animals, housing conditions, diet, etc.
- 364 Test conditions:
- the source, lot number, and manufacturer's quality assurance/quality control
 data (antibody sensitivity and specificity and the limit of detection) for the
- 367 ELISA kit;
- details of test substance preparation and application;
- justification for dose selection (including results from pre-screen test, if conducted);
- vehicle and test substance concentrations used, and total amount of test
 substance applied;
- details of food and water quality (including diet type/source, water source);
- details of treatment and sampling schedules;
- methods for measurement of toxicity;
- criteria for considering studies as positive or negative;

377 378	-	details of any protocol deviations and an explanation on how the deviation affects the study design and results.
379	Reliability che	eck:
380 381 382 383 384 385 386	- - -	a summary of results of latest reliability check, including information on test substance, concentration and vehicle used; concurrent and/or historical PC and concurrent negative (solvent/vehicle) control data for testing laboratory; if a concurrent PC was not included, the date and laboratory report for the most recent periodic PC and a report detailing the historical PC data for the laboratory justifying the basis for not conducting a concurrent PC.
387	Results:	
388 389 390 391 392 393 394 395 396 397 398 399 400	- - - -	individual weights of mice at start of dosing and at scheduled kill; as well as mean and associated error term (e.g. SD, SEM) for each treatment group; time course of onset and signs of toxicity, including dermal irritation at site of administration, if any, for each animal; a table of individual mouse BrdU labelling indices and SI values for each treatment group; mean and associated error term (e.g. SD, SEM) for BrdU labelling index/mouse for each treatment group and the results of outlier analysis for each treatment group; calculated SI and an appropriate measure of variability that takes into account the inter-animal variability in both the test substance and control groups; dose response relationship; statistical analyses, where appropriate.
401	Discussion of	results:
402 403 404 405	-	a brief commentary on the results, the dose-response analysis, and statistical analyses, where appropriate, with a conclusion as to whether the test substance should be considered a skin sensitizer.

406 **LITERATURE**

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522	ANNEX 1
523	DEFINITIONS
524 525 526 527	Accuracy: The closeness of agreement between test method results and accepted reference values. It is a measure of test method performance and one aspect of relevance. The term is often used interchangeably with "concordance" to mean the proportion of correct outcomes of a test method.
528 529 530 531 532	Benchmark test substance: A sensitizing or non-sensitizing substance used as a standard for comparison to a test substance. A benchmark substance should have the following properties; (i) a consistent and reliable source(s); (ii) structural and functional similarity to the class of substances being tested; (iii) known physical/chemical characteristics; (iv) supporting data on known effects, and (v) known potency in the range of the desired response.
533 534	False negative: A test substance incorrectly identified as negative or non-active by a test method, when in fact it is positive or active.
535 536	False positive: A test substance incorrectly identified as positive or active by a test, when in fact it is negative or non-active.
537 538	Hazard: The potential for an adverse health or ecological effect. The adverse effect is manifested only if there is an exposure of sufficient level.
539 540 541 542 543 544	Inter-laboratory reproducibility: A measure of the extent to which different qualified laboratories, using the same protocol and testing the same test substance, can produce qualitatively and quantitatively similar results. Inter-laboratory reproducibility is determined during the prevalidation and validation processes, and indicates the extent to which a test can be successfully transferred between laboratories, also referred to as between-laboratory reproducibility.
545 546 547	Intra-laboratory reproducibility: A determination of the extent that qualified people within the same laboratory can successfully replicate results using a specific protocol at different times. Also referred to as within-laboratory reproducibility.
548 549	Outlier: An outlier is an observation that is markedly different from other values in a random sample from a population.
550 551 552	Quality assurance: A management process by which adherence to laboratory testing standards, requirements, and record keeping procedures, and the accuracy of data transfer, are assessed by individuals who are independent from those performing the testing.
553554555	Reliability: Measures of the extent that a test method can be performed reproducibly within and between laboratories over time, when performed using the same protocol. It is assessed by calculating intra- and inter-laboratory reproducibility.
556557558	Skin sensitization: An immunological process that results when a susceptible individual is exposed topically to an inducing chemical allergen, which provokes a cutaneous immune response that can lead to the development of contact sensitization.
559 560 561	Stimulation Index (SI): A value calculated to assess the skin sensitization potential of a test substance that is the ratio of the proliferation in treated groups to that in the concurrent vehicle control group.

Test substance: Any material tested using this TG, whether it is a single compound or consists of multiple components (*e.g.* final products, formulations). When testing formulations, consideration should be given to the fact that certain regulatory authorities only require testing of the final product formulation. However, there may also be testing requirements for the active ingredient(s) of a product formulation.